

08/945144
41 Rec'd PCT/PTO 14 OCT 1997

MUTATED 5-ENOL
PYRUVYLSHIKIMATE-3-PHOSPHATE SYNTHASE,
GENE CODING
FOR SAID PROTEIN
AND TRANSFORMED PLANTS
CONTAINING SAID GENE

Michel Lebrun
Alain Sailland
Georges Freyssinet
and
Eric DeGryse

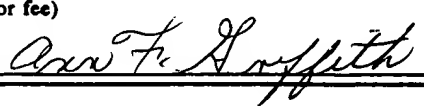
INTERNATIONAL APPLICATION
IN ENGLISH
including
SEARCH REPORT

RP/PCT

PCT/FR96/01125

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IFD: 07/18/96

<p>"Express Mail" mailing label number <u>ET841020925 IIS</u></p>
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<p><u>ANN F. GRIFFITH</u> (Typed or Printed name of person mailing paper or fee)</p>


Mutated 5-enolpyruvylshikimate-3-phosphate synthase,
gene coding for this protein and transformed plants
containing this gene

C + SU+CL \rightarrow Field of the Invention

The present invention relates to a new

5 5-enolpyruvylshikimate-3-phosphate synthase (or EPSPS)
which displays increased tolerance with respect to
herbicides which are competitive inhibitors with
respect to phosphoenolpyruvate (PEP) of EPSPS activity.
This more tolerant EPSPS synthase possesses at least one
10 "threonine by isoleucine" substitution. The invention
also relates to a gene coding for such a protein, to
plant cells transformed by chimeric gene constructions
containing this gene, to the plants regenerated from
these cells and also to the plants originating from
15 crossing using these transformed plants.

CL

\rightarrow Background of the Invention

Glyphosate, sulfosate and fosametine are
broad-spectrum systemic herbicides of the
phosphonomethylglycine family. They act essentially as
competitive inhibitors of 5-enolpyruvylshikimate-3-
20 phosphate synthase (EC 2.5.1.19) or EPSPS with respect
to the PEP (phosphoenolpyruvate). After their
application to the plant, they are translocated in the
plant where they accumulate in the rapidly growing
parts, in particular the cauline and root apices,
25 causing damage to the point of destruction of sensitive
plants.

Plastid EPSPS, the main target of these

2

products, is an enzyme of the pathway of biosynthesis of aromatic amino acids, which is encoded by one or more nuclear genes and synthesized in the form of a cytoplasmic precursor, then imported into the plastids 5 where it accumulates in its mature form.

The tolerance of plants to glyphosate and to products of the family is obtained by stable introduction into their genome of an EPSPS gene, of plant or bacterial origin, which is mutated or 10 otherwise in respect of the characteristics of inhibition by glyphosate of the product of this gene. In view of the mode of action of glyphosate and the degree of tolerance to glyphosate of the product of the genes which are used, it is advantageous to be able to 15 express the product of the translation of this gene so as enable it to be accumulated in substantial amounts in the plastids.

It is known, for example from US Patent 4,535,060, to confer on a plant a tolerance to a 20 herbicide of the above type, especially N-phosphono-methylglycine or glyphosate, by introducing into the genome of plants a gene coding for an EPSPS carrying at least one mutation that makes this enzyme more resistant to its competitive inhibitor (glyphosate) 25 after localization of the enzyme in the plastid compartment. These techniques, however, need to be improved in order to obtain greater reliability in the use of these plants under agricultural conditions.

CL *Summary of the Invention*³

In the present description, "plant" is understood to mean any differentiated multicellular organism capable of photosynthesis, and "plant cell" is understood to mean any cell originating from a plant
5 and capable of constituting undifferentiated tissues such as calluses or differentiated tissues such as embryos or plant parts or seeds.

The subject of the present invention is the production of transformed plants having increased
10 tolerance to herbicides of the phosphonomethylglycine family, by regeneration of cells transformed by means of new chimeric genes containing a gene for tolerance to these herbicides.

The subject of the invention is also a
15 chimeric gene for conferring on plants increased tolerance with respect to a herbicide having EPSPS as its target, comprising, in the direction of transcription: a promoter region, optionally a transit peptide region, a sequence of a gene coding for a
20 glyphosate tolerance enzyme and an untranslated polyadenylation signal region at the 3' end, characterized in that the glyphosate tolerance gene contains, relative to the gene from which it is derived, a "threonine 102 by isoleucine" substitution
25 in the "aroA" (EPSPS) region. Preferably, it comprises, in addition, in the same region, a "proline 106 by serine" substitution. These substitutions can be introduced or be present in an EPSPS sequence of any

origin, in particular of plant, bacterial, algal or fungal origin.

CL 1 Description of Preferred Embodiments

The transit peptides which can be used in the transit peptide region can be, known per se, of plant origin, for example originating from maize, sunflower, pea, tobacco or the like. The first and the second transit peptide can be identical, similar or different. They can, in addition, each comprise one or more transit peptide units according to European Patent 10 Application EP 0 508 909. It is the role of this characteristic region to permit the release of a mature and native protein, and especially the above mutated EPSPS, with maximum efficacy in the plasmid compartment.

15 The promoter region of the chimeric gene according to the invention may be advantageously composed of at least one gene promoter or promoter fragment which is expressed naturally in plants (tubulin, introns, actin, histone).

20 The untranslated transcription termination signal region at the 3' end of the chimeric gene may be of any origin, for example of bacterial origin, such as that of the nopaline synthase gene, or of plant origin, such as that of the *Arabidopsis thaliana* histone H4A748 25 gene according to the European Patent Application (European Application 633 317).

The chimeric gene according to the invention can comprise, in addition to the essential portions

above, at least one untranslated intermediate (linker) region, which can be located between the different transcribed regions described above. This intermediate region can be of any origin, for example of bacterial, 5 viral or plant origin.

Isolation of a cDNA coding for a maize EPSPS:

DE P The different steps which led to the obtaining of maize EPSPS cDNA, which served as substrate for the introduction of the two mutations, 10 are described below. All the operations described below are given by way of example, and correspond to a choice made from among the different methods available for arriving at the same result. This choice has no effect on the quality of the result, and consequently any 15 suitable method may be used by a person skilled in the art to arrive at the same result. Most of the methods of engineering of DNA fragments are described in "Current Protocols in Molecular Biology" Volumes 1 and 2, Ausubel F.M. et al., published by Greene Publishing 20 Associates and Wiley-Interscience (1989) (hereinafter, references to protocols described in this work will be designated "ref. CPMB"). The operations relating to DNA which were performed according to the protocols described in this work are especially the following: 25 ligation of DNA fragments, treatment with Klenow DNA polymerase and T4 DNA polymerase, preparation of plasmid and of bacteriophage λ DNA, either as a minipreparation or as a maxipreparation, and DNA and

RNA analyses according to the Southern and Northern techniques, respectively. Other methods described in this work were followed, and only significant modifications or additions to these protocols have been 5 described below.

CLV/C

Example 1:

P1 1. Obtaining of an *Arabidopsis thaliana* EPSPS fragment

10 P1 a) Two 20-mer oligonucleotides of respective sequences:

C L 5'-GCTCTGCTCATGTCTGCTCC-3'
 CL 5'-GCCCGCCCTTGACAAAGAAA-3'

(SEQ ID NO:6)

(SEQ ID NO:7)

15 were synthesized from the sequence of an *Arabidopsis thaliana* EPSPS gene (Klee H.J. et al. (1987) Mol. Gen. Genet., 210, 437-442). These two oligonucleotides are at positions 1523 to 1543 and 1737 to 1717, respectively, of the published sequence, and in opposite orientations.

20 P1 b) *Arabidopsis thaliana* (var. *columbia*) total DNA was obtained from Clontech (catalogue reference: 6970-1).

25 P1 c) 50 nanograms (ng) of DNA are mixed with 300 ng of each of the oligonucleotides and subjected to 35 amplification cycles with a Perkin-Elmer 9600 apparatus, under the conditions of standard medium for amplification which are recommended by the supplier.

The resulting 204-bp fragment constitutes the

Arabidopsis thaliana EPSPS fragment.

P1

2. Construction of a library of a cDNA from a
BMS maize cell line

P1 a) 5 g of filtered cells are ground in liquid

5 nitrogen, and the total nucleic acids are extracted
according to the method described by Shure et al. with
the following modifications:

P1 - the pH of the lysis buffer is adjusted
to pH 9.0;

10 P1 - after precipitation with isopropanol,
the pellet is taken up in water and,
after dissolution, adjusted to 2.5 M
LiCl. After incubation for 12 h at °C,
the pellet from centrifugation for 15
15 min at 30,000 g at 4°C is
resolubilized. The LiCl precipitation
step is then repeated. The resolubilized
pellet constitutes the RNA fraction of
the total nucleic acids.

20 P1 b) The poly(A)⁺ RNA fraction of the RNA
fraction is obtained by chromatography on an oligo(dT)-
cellulose column as described in "Current Protocols in
Molecular Biology".

P1 c) Synthesis of double-stranded cDNA having a
25 synthetic EcoRI end: this is carried out according to
the protocol of the supplier of the different reagents
needed for this synthesis in the form of a kit: the
"copy kit" from the company In Vitrogen.

Two single-stranded and partially complementary oligonucleotides of respective sequences:

CL 5'-AATTCCCCGGG-3'

CL 5'-CCCGGG-3' (the latter being
5 phosphorylated)

are ligated with the blunt-ended double-stranded cDNAs.

P This ligation of the adaptors results in the creation of SmaI sites attached to the double-stranded cDNAs and EcoRI sites in cohesive form at each end of
10 the double-stranded cDNAs.

P d) Creation of the library:

P The cDNAs possessing the artificial cohesive EcoRI sites at their ends are ligated with bacteriophage λgt10 cDNA which has been cut with EcoRI
15 and dephosphorylated according to the protocol of the supplier New England Biolabs.

P An aliquot of the ligation reaction was encapsidated in vitro with encapsidation extracts, namely Gigapack Gold, according to the supplier's
20 instructions; this library was titrated using the bacterium *E. coli* C600hfl. The library thereby obtained is amplified and stored according to the instructions of the same supplier, and constitutes the BMS maize cell suspension cDNA library.

25 P+b 3. Screening of the BMS maize cell suspension cDNA library with the *Arabidopsis thaliana* EPSP probe

P The protocol followed is that of "Current Protocols in Molecular Biology" Volumes 1 and 2,

Ausubel F.M. et al., published by Greene Publishing
Associates and Wiley-Interscience (1989) (CPMB).
Briefly, approximately 10^6 recombinant phages are
plated out on LB dishes at an average density of 100
5 phages/cm². The lytic plaques are replicated in
duplicate on Amersham Hybond N membranes.

P The DNA was fixed to the filters by 1600kJ UV
treatment (Stratagene Stratalinker). The filters were
prehybridized in 6xSSC/0.1%SDS/0.25 skinned milk for
10 2 h at 65°C. The *Arabidopsis thaliana* EPSPS probe was
labelled with [³²P]dCTP by random priming according to
the supplier's instructions (Pharmacia Ready to Go
kit). The specific activity obtained is of the order of
15 10^8 cpm per μ g of fragment. After denaturation for 5
min at 100°C, the probe is added to the
prehybridization medium and hybridization is continued
for 14 hours at 55°C. The filters are fluorographed for
48 h at -80°C with Kodak XAR5 film and Amersham
Hyperscreen RPN enhancing screens. Alignment of the
20 positive spots on the filter with the dishes from which
they originate enables zones corresponding to the
phages displaying a positive hybridization response
with the *Arabidopsis thaliana* EPSPS probe to be picked
out from the dish. This step of plating out, transfer,
25 hybridization and recovery is repeated until all the
spots in the dish of the successively purified phages
prove 100% positive in hybridization. An independent
plaque of phage lysis is then picked out in diluent λ

10

medium (Tris-Cl pH 7.5; 10mM MgSO₄; 0.1M NaCl; 0.1% gelatin); these phages in solution constitute the EPSPS-positive clones of the BMS maize cell suspension.

Pt b 4. Preparation and analysis of the DNA of the
5 EPSPS clones of the BMS maize cell suspension

P Approximately 5×10⁸ phages are added to 20 ml of C600hfl bacteria at an OD_{600nm} value of 2/ml and incubated for 15 minutes at 37°C. This suspension is then diluted in 200 ml of bacterial growth medium in a
10 1-l Erlenmeyer and stirred in a rotary stirrer at 250 rpm. Lysis is noted when the medium clarifies, corresponding to the lysis of the turbid bacteria, and takes place after approximately 4 h of stirring. This supernatant is then treated as described in "Current
15 Protocols in Molecular Biology". The DNA obtained corresponds to the EPSPS clones of the BMS maize cell suspension.

One to two µg of this DNA are cut with EcoRI and separated on 0.8% LGTA/TBE agarose gel (ref. CPMB).
20 A final verification consists in checking that the purified DNA does indeed display a hybridization signal with the *Arabidopsis thaliana* EPSPS probe. After electrophoresis, the DNA fragments are transferred onto Amersham Hybond N membranes according to the protocol
25 of Southern described in "Current Protocols in Molecular Biology". The filter is hybridized with the *Arabidopsis thaliana* EPSPS probe according to the conditions described in section 3 above. The clone

displaying a hybridization signal with the *Arabidopsis thaliana* EPSPS probe and containing the longest EcoRI fragment has a size estimated on gel as approximately 1.7 kbp.

5 *Pf6* 5. Obtaining of the clone pRPA-ML-711
P Ten μ g of the phage clone containing the 1.7-kbp insert are digested with EcoRI and separated on 0.8% LGTA/TBE agarose gel (ref. CPMB). The gel fragment containing the 1.7-kbp insert is excised from the gel
10 by BBT staining, and the fragment is treated with β -agarase according to the protocol of the supplier, New England Biolabs. The purified DNA of the 1.7-kbp fragment is ligated at 12°C for 14 h with the DNA of plasmid pUC 19 (New England Biolabs) cut with EcoRI
15 according to the ligation protocol described in "Current Protocols in Molecular Biology". Two μ l of the above ligation mixture are used for the transformation of an aliquot of electrocompetent *E. coli* DH10B; transformation is accomplished by electroporation using
20 the following conditions: the mixture of competent bacteria and ligation medium is introduced into an electroporation cell of thickness 0.2 cm (Biorad) previously cooled to 0°C. The physical conditions of the electroporation using an electroporator made by
25 Biorad are 2500 volts, 25 μ F and 200 Ω . Under these conditions, the mean discharge time of the condenser is of the order of 4.2 milliseconds. The bacteria are then taken up in 1 ml of SOC medium (ref. CPMB) and stirred

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for 1 hour at 200 rpm on a rotary stirrer in 15-ml Corning tubes. After plating out on LB/agar medium supplemented with 100 μ g/ml of carbenicillin, minipreparations of the bacterial clones which have grown

5 after one night at 37°C are produced according to the protocol described in "Current Protocols in Molecular Biology". After digestion of the DNA with EcoRI and separation by electrophoresis on 0.8% LGTA/TBE agarose gel (ref. CPMB), the clones possessing a 1.7-kbp insert

10 are retained. A final verification consists in checking that the purified DNA does indeed display a hybridization signal with the *Arabidopsis thaliana* EPSPS probe. After electrophoresis, the DNA fragments are transferred onto Amersham Hybond N membranes

15 according to the protocol of Southern described in "Current Protocols in Molecular Biology". The filter is hybridized with the *Arabidopsis thaliana* EPSPS probe according to the conditions described in section 3 above. The plasmid clone possessing a 1.7-kbp insert

20 and hybridizing with the *Arabidopsis thaliana* EPSPS probe was prepared on a larger scale, and the DNA resulting from the lysis of the bacteria was purified on a CsCl gradient as described in "Current Protocols in Molecular Biology". The purified DNA was partially

25 sequenced with a Pharmacia kit according to the supplier's instructions and using as primers the M13 direct and reverse universal primers ordered from the same supplier. The partial sequence produced covers

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approximately 0.5 kbp. The derived amino acid sequence in the region of the mature protein (approximately 50 amino acid residues) displays 100% identity with the corresponding amino sequence of mature maize EPSPS

5 described in American Patent USP 4,971,908. This clone, corresponding to a 1.7-kbp EcoRI fragment of the EPSP DNA of the BMS maize cell suspension, was designated pRPA-ML-711. The complete sequence of this clone was determined on both strands using the protocol of the

10 Pharmacia kit and synthesizing complementary oligonucleotides and those of the opposite orientation every 250 bp approximately. The complete sequence obtained of this 1713-bp clone is presented in SEQ ID No. 1.

15 *Pt b* 6. Obtaining of the clone pRPA-ML-715

P Analysis of the sequence of the clone pRPA-ML-711, and especially comparison of the derived amino acid sequence with that of maize, shows a sequence extension of 92 bp upstream of the GCG codon coding for the NH₂-terminal alanine of the mature portion of maize EPSPS (American Patent USP 4,971,908). Similarly, an extension of 288 bp downstream of the AAT codon coding for the COOH-terminal asparagine of the mature portion of maize EPSPS (American Patent USP 4,971,908) is observed. These two portions could correspond, in the case of the NH₂-terminal extension to a portion of the sequence of a transit peptide for plastid localization, and, in the case of the COOH-terminal extension, to the

untranslated 3' region of the cDNA.

P In order to obtain a cDNA coding for the mature portion of the maize EPSPS cDNA, as described in USP 4,971,908, the following operations were carried
5 out:

P a) Removal of the untranslated 3' region:
construction of pRPA-ML-712:

P The clone pRPA-ML-711 was cut with the restriction enzyme AseI, and the ends resulting from
10 this cleavage were rendered blunt by treatment with the Klenow fragment of DNA polymerase I according to the protocol described in CPMB. A cleavage with the restriction enzyme SacII was then performed. The DNA resulting from these operations was separated by
15 electrophoresis on 1% LGTA/TBE agarose gel (ref. CPMB).

P The gel fragment containing the 0.4-kbp "AseI-blunt ends/SacII" insert was excised from the gel and purified according to the protocol described in section 5 above. The DNA of the clone pRPA-ML-711 was
20 cut with the restriction enzyme HindIII at the HindIII site located in the polylinker of the cloning vector pUC19, and the ends resulting from this cleavage were rendered blunt by treatment with the Klenow fragment of DNA polymerase I. A cleavage with the restriction
25 enzyme SacII was then performed. The DNA resulting from these manipulations was separated by electrophoresis on 0.7% LGTA/TBE agarose gel (ref. CPMB).

P The gel fragment containing the approximately

3.7-kbp HindIII-blunt ends/SacII insert was excised from the gel and purified according to the protocol described in section 5 above.

P The two inserts were ligated, and 2 μ l of the 5 ligation mixture were used to transform *E. coli* DH10B as described above in section 5.

P The plasmid DNA content of different clones was analysed according to the procedure described for pRPA-ML-711. One of the plasmid clones selected 10 contains an approximately 1.45-kbp EcoRI-HindIII insert. The sequence of the terminal ends of this clone reveals that the 5' end of the insert corresponds exactly to the corresponding end of pRPA-ML-711, and that the 3'-terminal end possesses the following 15 sequence:

CL "5' - ... AATTAAGCTCTAGAGTCGACCTGCAGGCATGCAAGCTT - 3'".

(SEQ ID NO: 8)

P The underlined sequence corresponds to the codon of the COOH-terminal amino acid asparagine, the next codon corresponding to the translation stop codon. 20 The nucleotides downstream correspond to sequence elements of the pUC19 polylinker. This clone comprising the pRPA-ML-711 sequence up to the translation termination site of mature maize EPSPS and followed by sequences of the pUC 19 polylinker up to the HindIII site was designated pRPA-ML-712.

P b) Modification of the 5' end of pRPA-ML-712: construction of pRPA-ML-715:

P The clone pRPA-ML-712 was cut with the

restriction enzymes *Pst*I and *Hind*III. The DNA resulting from these manipulations was separated by electrophoresis on 0.8% *LGTA/TBE* agarose gel (ref. CPMB). The gel fragment containing the 1.3-kbp *Pst*I-

5 *Eco*RI insert was excised from the gel and purified according to the protocol described in section 5 above. This insert was ligated in the presence of an equimolecular amount of each of the two partially complementary oligonucleotides of sequence:

CL 10 Oligo 1: 5'-GAGCCGAGCTCCATGGCCGGCGCCGAGGAGATCGTGCTGCA-3'
CL Oligo 2: 5'-GCACGATCTCCTCGCGCCGCCATGGAGCTCGGCTC-3' (SEQ ID NO:10)
PS as well as in the presence of plasmid pUC19 DNA digested with the restriction enzymes *Bam*HI and *Hind*III.

15 P Two μ l of the ligation mixture were used to transform *E. coli* DH10B as described above in section 5. After analysis of the plasmid DNA content of different clones according to the procedure described above in section 5, one of the clones possessing an 20 approximately 1.3-kbp insert was retained for subsequent analyses. The sequence of the 5'-terminal end of the selected clone reveals that the DNA sequence in this region is the following: sequence of the pUC19 polylinker from the *Eco*RI to the *Bam*HI sites, followed 25 by the sequence of the oligonucleotides used in the cloning, followed by the remainder of the sequence present in pRPA-ML-712. This clone was designated pRPA-ML-713. This clone possesses a methionine ATG codon

included in an NcoI site upstream of the N-terminal alanine codon of mature EPSP synthase. Furthermore, the alanine and glycine codons of the N-terminal end have been preserved, but modified on the third variable 5 base: initial GCGGGT gives modified GCGGGC.

P The clone pRPA-ML-713 was cut with the restriction enzyme HindIII, and the ends of this cleavage were rendered blunt by treatment with the Klenow fragment of DNA polymerase I. A cleavage with 10 restriction enzyme SacI was then performed. The DNA resulting from these manipulations was separated by electrophoresis on 0.8% LGTA/TBE agarose gel (ref. CPMB). The gel fragment containing the 1.3-kbp "HindIII-blunt ends/SacI" insert was excised from the 15 gel and purified according to the protocol described in section 5 above. This insert was ligated in the presence of plasmid pUC19 DNA digested with restriction enzyme XbaI, and the ends of this cleavage were rendered blunt by treatment with the Klenow fragment of 20 DNA polymerase I. A cleavage with the restriction enzyme SacI was then performed. Two μ l of the ligation mixture were used to transform *E. coli* DH10B as described above in section 5. After analysis of the plasmid DNA content of different clones according to 25 the procedure described above in section 5, one of the clones possessing an approximately 1.3-kbp insert was retained for subsequent analyses. The sequence of the terminal ends of the selected clone reveals that the

DNA sequence is the following: sequence of the pUC19 polylinker from the EcoRI to SacI sites, followed by the sequence of the oligonucleotides used in the cloning from which the 4 bp GATCC of the

5 oligonucleotide 1 described above have been deleted, followed by the remainder of the sequence present in pRPA-ML-712 up to the HindIII site and sequence of the pUC19 polylinker from XbaI to HindIII. This clone was designated pRPA-ML-715.

10 *Pf6* 7. Obtaining of a cDNA coding for a mutated maize EPSPS

P All the mutagenesis steps were carried out with the Pharmacia U.S.E. mutagenesis kit according to the supplier's instructions. The principle of this

15 mutagenesis system is as follows: plasmid DNA is denatured by heat and reassociated in the presence of a molar excess of, on the one hand the mutagenesis oligonucleotide, and on the other hand an oligonucleotide enabling a unique restriction enzyme site present in

20 the polylinker to be eliminated. After the reassociation step, synthesis of the complementary strand is carried out by the action of T4 DNA polymerase in the presence of T4 DNA ligase and gene 32 protein in a suitable buffer which is supplied. The

25 synthesis product is incubated in the presence of the restriction enzyme for which the site is assumed to have disappeared by mutagenesis. The *E. coli* strain possessing, in particular, the mutS mutation is used as

host for the transformation of this DNA. After growth in liquid medium, the total plasmid DNA is prepared and incubated in the presence of the restriction enzyme used before. After these treatments, *E. coli* strain 5 DH10B is used as host for the transformation. The plasmid DNA of the clones isolated is prepared, and the presence of the mutation introduced is verified by sequencing.

A) - modification of sites or sequences

10 without in principle affecting the EPSPS-resistance character of maize to products which are competitive inhibitors of EPSP synthase activity: elimination of an internal NcoI site from pRPA-ML-715.

15 The pRPA-ML-715 sequence is numbered arbitrarily by placing the first base of the N-terminal alanine codon GCC at position 1. This sequence possesses an NcoI site at position 1217. The site-modification oligonucleotide possesses the sequence:

CL 5' -CCACAGGATGGCGATGGCCTTCTCC-3' ^(SEQ ID NO:11)

20 P After sequencing according to the references given above, the sequence read after mutagenesis corresponds to that of the oligonucleotide used. The NcoI site has indeed been eliminated, and the translation into amino acids in this region preserves the initial 25 sequence present in pRPA-ML-715.

P This clone was designated pRPA-ML-716.

P The 1340-bp sequence of this clone is presented in SEQ ID No. 2 and SEQ ID No. 3.

20

P1

B)- sequence modifications enabling the EPSPS-resistance character of maize to products which are competitive inhibitors of EPSP synthase activity to be increased.

5

P The following oligonucleotides were used:

P1

a) mutation Thr 102 → Ile.

(SEQ ID NO: 13)

CL

5' -GAATGCTGGAATCGCAATGCGGCCATTGACAGC-3'

P1

b) mutation Pro 106 → Ser.

(SEQ ID NO: 13)

CL

5' -GAATGCTGGAATGCAATGCGGTCTTGACAGC-3'

P1

10

c) mutations Gly 101 → Ala and Thr 102 → Ile.

CL

5' -CTTGGGAAATGCTGCCATCGCAATGCGGCCATTG-3'

(SEQ ID NO: 14)

P1

d) mutations Thr 102 → Ile and Pro 106 → Ser.

(SEQ ID NO: 15)

CL

5' -GGGAAATGCTGGAATCGCAATGCGGTCTTGACAGC-3'

1

15

P After sequencing, the sequence read after

15

mutagenesis on the three mutated fragments is identical

to the parent pRPA-ML-716 DNA sequence, with the

exception of the mutagenized region which corresponds

to that of the mutagenesis oligonucleotides used. These

clones were designated: pRPA-ML-717 for the mutation

20

Thr 102 → Ile, pRPA-ML-718 for the mutation Pro 106 →

Ser, pRPA-ML-719 for the mutations Gly 101 → Ala and

Thr 102 → Ile and pRPA-ML-720 for the mutations Thr 102

→ Ile and Pro 106 → Ser.

P The 1340-bp sequence of pRPA-ML-720 is

presented in SEQ ID No. 4 and SEQ ID No. 5.

The 1395-bp NcoI-HindIII insert is the basis of all the constructions used for the transformation of plants for the introduction of resistance to herbicides which are competitive inhibitors of EPSPS, and especially glyphosate resistance. This insert will be designated in the remainder of the description "the maize EPSPS double mutant".

CLU/C Example 2:

10 CLUL Glyphosate tolerance of the different mutants
in vitro

P + b 2.a: Extraction of EPSP synthase

The different EPSP synthase genes are introduced in the form of an NcoI-HindIII cassette into 15 the plasmid vector pTrc99a (Pharmacia, ref: 27-5007-01) cut with NcoI and HindIII. Recombinant *E. coli* DH10B bacteria overexpressing the different EPSP synthases are sonicated in 40 ml of buffer per 10 g of pelleted cells, and washed with this same buffer (200 mM Tris-HCl pH 7.8, 50 mM mercaptoethanol, 5 mM EDTA and 1 mM PMSF), to which 1 g of polyvinylpyrrolidone is added. The suspension is stirred for 15 minutes at 4°C and then centrifuged for 20 minutes at 27,000 g and 4°C.

Ammonium sulphate is added to the supernatant 25 to bring the solution to 40% saturation with respect to ammonium sulphate. The mixture is centrifuged for 20 minutes at 27,000 g and 4°C. Ammonium sulphate is added to the new supernatant to bring the solution to

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70% saturation with respect to ammonium sulphate. The mixture is centrifuged for 30 minutes at 27,000 g and 4°C. The EPSP synthase present in this protein pellet is taken up in 1 ml of buffer (20 mM Tris-HCl pH 7.8 and 50 mM mercaptoethanol). This solution is dialysed overnight against two litres of this same buffer at 4°C.

P+b 2.b: Enzyme activity

The activity of each enzyme, as well as its 10 glyphosate resistance, is measured *in vitro* over 10 minutes at 37°C in the following reaction mixture: 100 mM maleic acid pH 5.6, 1 mM phosphoenolpyruvate, 3 mM shikimate 3-phosphate (prepared according to Knowles P.F. and Sprinson D.B. 1970. Methods in Enzymol 15 17A, 351-352 from *Aerobacter aerogenes* strain ATCC 25597) and 10 mM potassium fluoride. The enzyme extract is added at the last moment after the addition of glyphosate, the final concentration of which varies from 0 to 20 mM.

20 *P* The activity is measured by assaying the phosphate liberated according to the technique of Tausky H.A. and Shorr E. 1953. *J. Biol. Chem.* 202, 675-685.

25 *P* Under these conditions, the wild-type (WT) enzyme is already 85% inhibited at a glyphosate concentration of 0.12 mM. At this concentration, the mutant enzyme known as Ser106 is only 50% inhibited, and the other three mutants, Ile102, Ile102/Ser106 and

Ala101/Ile102, show little or no inhibition.

The glyphosate concentration has to be multiplied by ten, that is to say 1.2 mM, in order to produce a 50% inhibition of the mutant enzyme Ile102.

5 the mutants Ile102/Ser106, Ala/Ile and Ala still not being inhibited.

It should be noted that the activity of the mutants Ala/Ile and Ala is not inhibited up to glyphosate concentrations of 10mM, and that that of the 10 mutant Ile102/Ser106 is not reduced even if the glyphosate concentration is multiplied by 2, that is to say 20 mM.

CL U/C Example 3:

CL U/L Resistance of transformed tobacco plants

15 P 1-1- Transformation

P The vector pRPA-RD-173 is introduced into *Agrobacterium tumefaciens* strain EHA101 (Hood et al., 1987) carrying the cosmid pTVK291 (Komari et al., 1986). The transformation technique is based on the 20 procedure of Horsh et al. (1985).

P 1-2- Regeneration

P The regeneration of PBD6 tobacco (source SEITA France) from leaf explants is carried out on a Murashige and Skoog (MS) basal medium comprising 30 g/l 25 of sucrose as well as 200 µg/ml of kanamycin. The leaf explants are removed from plants cultivated in the greenhouse or in vitro and are transformed according to the leaf disc technique (Science, 1985, Vol. 227, pp.

1229-1231) in three successive steps: the first comprises the induction of shoots on a medium supplemented with 30 g/l of sucrose containing 0.05 mg/l of naphthylacetic acid (NAA) and 2 mg/l of 5 benzylaminopurine (BAP) for 15 days. The shoots formed during this step are then developed for 10 days by culturing on an MS medium supplemented with 30 g/l of sucrose but not containing any hormone. Shoots which have developed are then removed and cultured on an MS 10 rooting medium having half the content of salts, vitamins and sugar and not containing any hormone. After approximately 15 days, the rooted shoots are transferred to soil.

P 1-3- Glyphosate resistance

15 P Twenty transformed plants were regenerated and transferred to the greenhouse for the construction of pRPA-RD-173. These plants were treated in the greenhouse at the 5-leaf stage with an aqueous suspension of RoundUp corresponding to 0.8 kg of 20 glyphosate active substance per hectare.

The results correspond to the observation of phytotoxicity indices recorded 3 weeks after treatment. Under these conditions, it is found that the plants transformed with the construction pRPA-RD-173 display 25 very good tolerance, whereas the untransformed control plants are completely destroyed.

These results show clearly the improvement brought about by the use of a chimeric gene according

to the invention for the same gene coding for
glyphosate tolerance.

CL VI/C Example 4:

CL VI/L Transformation and selection of maize cells

5 P BMS (Black Mexican Sweet) maize cells in an
exponential growth phase are bombarded with the
construction pRPA-RD-130 according to the principle and
the protocol described by Klein et al. 1987 (Klein
T.M., Wolf E.D., Wu R. and Sandford J.C. (1987): High
10 velocity microprojectiles for delivering nucleic acids
into living cells, NATURE Vol. 327 pp. 70-73).

P Two days after bombardment, the cells are
transferred to the same medium containing 2 mM
N-(phosphonomethyl)glycine.

15 After 8 weeks of selection on this medium,
calluses which develop are selected, then amplified and
analysed by PCR, and reveal clearly the presence of the
chimeric OTR-EPSPS gene.

Cells not bombarded and grown on the same
20 medium containing 2 mM N-(phosphonomethyl)glycine are
blocked by the herbicide and do not develop.

The transformed plants according to the
invention may be used as parents for obtaining lines
and hybrids having the phenotypic character
25 corresponding to the expression of the chimeric gene
introduced.

P Description of the constructions of the
plasmids

P pRPA-RD-124: Addition of a "nos"
polyadenylation signal to pRPA-ML-720 with creation of
5 a cloning cassette containing the maize double mutant
EPSPS gene (Thr 102 → Ile and Pro 106 → Ser). pRPA-ML-
720 is digested with HindIII and treated with the
Klenow fragment of *E. coli* DNA polymerase I to produce
a blunt end. A second digestion is performed with NcoI,
10 and the EPSPS fragment is purified. The EPSPS gene is
then ligated with purified pRPA-RD-12 (a cloning
cassette containing the polyadenylation signal of
nopaline synthase) to give pRPA-RD-124. To obtain the
useful purified vector pRPA-RD-12, it was necessary for
15 the latter to be digested beforehand with Sall, treated
with Klenow DNA polymerase and then digested a second
time with NcoI.

P pRPA-RD-125: Addition of an optimized transit
peptide (OTP) to pRPA-RD-124 with creation of a cloning
20 cassette containing the EPSPS gene targeted on the
plasmids. pRPA-RD-7 (European Patent Application
EP 652 286) is digested with SphI, treated with T4 DNA
polymerase and then digested with SpeI, and the OTP
fragment is purified. This OTP fragment is cloned into
25 pRPA-RD-124 which has previously been digested with
NcoI, treated with Klenow DNA polymerase to remove the
protruding 3' portion and then digested with SpeI. This
clone is then sequenced in order to ensure correct

translational fusion between the OTP and the EPSPS gene. pRPA-RD-125 is then obtained.

P pRPA-RD-130: Addition of the H3C4 maize histone promoter and of adhl intron 1 sequences of 5 pRPA-RD-123 (Patent Application EP 507 698) to pRPA-RD-125 with creation of a cassette for expression in plants for the expression of the double mutant EPSPS gene in the tissues of monocotyledons. pRPA-RD-123 (a cassette containing the H3C4 maize histone promoter 10 fused with the adhl intron 1) is digested with NcoI and SacI. The DNA fragment containing the promoter derived from pRPA-RD-123 is then purified and ligated with pRPA-RD-125 which has previously been digested with NcoI and SacI.

15 P pRPA-RD-159: Addition of the H4A748 *Arabidopsis* histone double promoter (Patent Application EP 507 698) to pRPA-RD-125 with creation of a cassette for expression in plants for the expression of the "OTP-double mutant EPSPS gene" gene in the tissues of 20 dicotyledons. pRPA-RD-132 (a cassette containing the H4A748 double promoter (Patent Application EP 507 698)) is digested with NcoI and SacI. The purified promoter fragment is then cloned into pRPA-RD-125 which has been digested with EcoI and SacI.

25 pRPA-RD-173: Addition of the "H4A748 promoter-OTP-double mutant EPSPS gene" gene of pRPA-RD-159 to plasmid pRPA-BL-150A (European Patent Application 508 909) with creation of an Agrobacterium

tumefaciens transformation vector. pRPA-RD-159 is digested with NotI and treated with Klenow polymerase. This fragment is then cloned into pRPA-BL-150A with SmaI.